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- (54) Nonaqueous-electrolyte secondary battery
  Sakundärbatterie mit nichtwässerigen Elektrolyten
  Batterie secondaire à électrolyte non-aqueux
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- (56) References cited; EP-A- 0 780 920 US-A- 5 508 122

US-A- 6 360 684

- PATENT ABSTRACTS OF JAPAN vol. 017, no. 583 (E-1477), 15 December 1993 & JP 05 234620 A (SONY CORP), 10 September 1993
- PATENT ABSTRACTS OF JAPAN vol. 012, no. 158 (E-808), 13 May 1988 & JP 62 272471 A (TOSHIBA CORP), 26 November 1987

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#### Description

### BACKGROUND OF THE INVENTION

#### Flatd of the Invention

(0001) The present invention relates to a nonequeous-electrolyte secondary battery incorporating a coll electrode formed by laminating elongated positive and negative electrodes through separators such that the outermost layer is the positive electrode.

## Related Background Art

[0002] Hitherto, secondary batterias for electronic apparatuses have been nickel-cadmium batterias or lead batteries. The progress of the electronic technique made in recent years has caused the performance of the electronic apparatus to be improved, the size to be reduced and a portable structure to be realized. As a result, a requirement has erisen to raise the energy density of the secondary battery for the electronic apparatus. However, there arises a problem in that the energy density of the nickel-cadmium battery and that of the lead battery cannot satisfactorily be relised because of low discharge voltages.

[0003] In recent years, a nonaqueous-electrolyte secondary battery has energetically been developed and researched as a secondary battery expected to be capable of raising the discharge voltage and resizing small salfdischarge and a long lifetime egalist cycle operations. The nonaqueous-electrolyte secondary battery has been employed in place of the nickel-cadmium battery and the lead battery. The nonaqueous-electrolyte secondary battery incorporates a negative electrode made of a meterial, such as a carbon material, which permits deping/dedoping tithium ions; and a positive electrode made of a composite lithium calde, such as composite lithium-cobalt outde.

[0004] As described above, the nonequeous-electrolyte secondary battery is required to have satisfactory charge-teristics including a discharge characteristic under a heavy load and lifetime against a cycle operation. Therefore, the structure of the electrodes of the above-mentioned nonequeous-electrolyte secondary battery has usually been formed into a coil electrode structure as shown in FIG. 1. As shown in FIG. 1, an elongated positive electrode 103 incorporates positive-electrode-mix tayers 102s and 102b formed by applying a positive-electrode ink to each of the two sides of a collector 101. An elongated negative electrode 108 similarly incorporates negative-electrode-mix tayers 105s and 105o formed by applying a negative-electrode mix to each of the two sides of a collector 104. The positive and negative electrodes 103 and 106 are wound such that a separator 107 is interposed so that a coll electrode 108 is formed, in the foregoing case, internal short circuit occurring when illhium is deposited during a charging operation must be prevented. Therefore, the width and length of the negative electrode 106 opposite to the positive electrode 103 usually are made to be larger than those of the positive electrode 103.

[0005] The above-mentioned coll electrode 108 incorporates the negative electrode 108 which forms the innermost layer and the outermost layer. Therefore, portions, each containing non-reacted negative-electrode active material which does not concern the charge/discharge, addt adjacent to the end of the outermost layer of the negative electrode 106 and the innermost layer of the same. Therefore, the inside portion of the battery cannot effectively be used. As a result, there arises a problem in that the energy density cannot satisfactorily be related.

[0006] To solve the above-mentioned problems, a technique has been disclosed in Japanese Patent Latd-Open No. 5-234620. As shown in FIG. 2, an elongated positive electrode 113 incorporates positive-electrode-mix layers 112a and 112b formed by applying a positive-electrode-mix to each of the two sides of a collector 111. An elongated negative electrode 116 incorporates negative-electrode-mix layers 115a and 115b formed by applying a negative-electrode-mix to each of the two sides of a collector 114. The positive electrode 113 and the negative electrode 116 are wound such that a separator 117 is interposed so that a coil electrode 118 is formed. This outermost layer electrode, with which charge/discharge of the coil electrode 118 is performed, is made to be the positive electrode 113. Moreover, a portion edjacent to an outermost and 113a of this positive electrode and/or a portion edjacent to an innermost end 113b is formed such that the positive-electrode-mix layer 112a in the foregoing case) is formed on only either material aurisos of the collector 111. Thus, the quantity of the non-reacted negative-electrode enterty can be reduced. Thus, this inside portion of the battery is effectively used to raise the energy density correspondingly.

[0007] However, the above-manifored coil electrode has the structure as shown in FIG. 2 such that the outermost and 116a of the negative electrode 116 is formed by only the collector 114. A negative-electrode lead 119 is provided for the upper surface of the collector 114 so that a projection is formed on the upper surface. Also the positive electrode 113 has an outermost end 113a formed by only the collector 111. If the projection exists as described above, the projection pleases the appearance 117 disposed between the negative electrode 116 and the positive electrode 119. Thus, the projection is undestrably brought into contact with the collector 111 of the positive electrode 113. It leads to

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a fact that short circuit easily takes place. As a result, a percent defective is raised and, therefore, reliability deteriorates.

#### SUMMARY OF THE INVENTION

[9006] In view of the foregoing, an object of the present invention is to provide a nanequeous-electrolyte secondary battery which reduce the quantity of a non-reacted active material for the negative electrode in the battery to effectively use the inside portion of the battery so as to raise the energy density, elongate the lifetime against a cycle operation, prevent a defect and improve the reliability.

[9009] According to one aspect of the present invention, there is provided a nonequeous-electrolyte secondary battery including: a coll electrode formed by terminating an elongated positive electrode which has a positive-electrode mix tayer formed on at least either of main surfaces of a positive-electrode collector and an etongated negative electrode which has a negative-electrode-mix tayer formed on at least either main surfaces of a negative-electrode collector and by whoding a formed laminate such that the positive electrode is positioned at the cultimost positive of the collector and wherein the positive-electrode-mix tayer is formed on only either of main surfaces of the collector at the positive electrode, wherein the positive electrode and/or the position adjacent to the imageness and of the positive electrode on the positive-electrode collector at the outermost and of the positive electrode end only the positive-electrode collector is formed, the negative-electrode collector at the outermost and of the negative-electrode and only the negative-electrode collector is formed, and the outermost and of the negative-electrode and only the negative-electrode collector is formed, and the outermost and of the negative-electrode to the outermost and of the negative-electrode collector positioned at the outermost and of the negative-electrode to the outermost and of the coll electrode toward the outer portion of the coll electrode collector.

[9010] The nonaqueous-electrolyte secondary battery according to the present invention may have a structure that the coil electrode incorporates a negative-electrode lead edjecent to an outermost end of the negative-electrode collector positioned at the outermost end of the negative-electrode, and the negative-electrode lead is positioned more forwards than the outermost end of the positive-electrode collector positioned at the outermost end of the positive electrode.

[0011] The nonequeous-electrolyte secondary battery according to the present invention may have a structure that the coil electrode has a structure that distance L from the cutermost end of the negative-electrode collector positioned at the outermost end of the negative-electrode collector positioned at the outermost end of the positive-electrode collector positioned at the outermost end of the positive electrode which are positioned in a fore-end-aft direction from the inner portion of the coil electrode toward the cuter portion of the coil electrode satisfies the following retailorship on an assumption that the diameter of the coil electrode is d:

## 0 < L ≤ #d

[0012] The nonaqueous-electrolyte secondary battery according to the present invention may have a structure that the negotive-electrode mix contains a negative-electrode material and a binder.

[0013] The nonaqueous-electrolyte secondary bettery according to the present invention may have a structure that the negative-electrode material is at least one type material selected from a group consisting of a crystalline metal oxide and an emorphous metal oxide which permit doping/dedoping lithium long.

[0014] The nonequeous-electrolyte secondary battery according to the present invention may have a structure that the positive-electrode mix contains a positive-electrode material, a conductive material and a binder.

(0015) The nonaqueous-electrolyte secondary battery according to the present invention may have a structure that the positive-electrode meterial is at least one type material selected from a group consisting of LIMO<sub>2</sub> (where M is at least any one of Co, NI, Min, Fe, AJ, V and TI) and interlayer compounds each containing LL.

[0016] The nonsqueous-electrolyte secondary battery according to the present invention may have a structure that the separator is made of at teast one type material selected from a group consisting of polyathylene and polypropytene. [0017] The nonsqueous-electrolyte secondary battery according to the present invention may have a structure that the positive-electrode collector is made of at least one type material selected from a group consisting of aluminum, statistical and nickel.

[0018] The nonequeous-electrolyte secondary battery according to the present invention may have a structure that the negative-electrode collector is made of at least one type material selected from a group consisting of copper, staintess steel and nickel.

[0019] The nonequeous-electrolyte secondary battery according to the present invention may have a structure that the nonequeous-electrolyte secondary battery contains a nonequeous electrolyte prepared by dissolving an electrolyte in nonequeous solvent is made of at least one type material selected from a group consisting of propylene carbonate, ethylene carbonate, 1, 2-directhoxyethane, 1, 2-dicthoxyethane, diethylenebonate,

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y-butyrolactone, tetrahydrofuran, 1,3-dioxolane, 4-methyl-1, 3-dioxolane, diethylether, sulfolane, methylsulfolane, scetimitale and propionitite.

[0020] The nonsqueous-electrolyte secondary bettery eccording to the present invention may have a structure that the electrolyte is at least one type material selected from a group consisting of LICIO<sub>4</sub>, LIA $_6$ , LIP $_6$ , LIB $_4$ , LIB $_6$ Ce $_6$ H $_6$ A $_6$ LICH, LIBY, LISO<sub>3</sub>CH<sub>3</sub> and LISO<sub>3</sub>CF<sub>3</sub>.

[0021] Other objects, features and advantages of the Invention will be evident from the following datalise description of the preferred embodiments described in conjunction with the attached drawings.

### BRIEF DESCRIPTION OF THE DRAWINGS

#### [0022]

FIG. 1 is a crose sectional view showing the structure of a conventional nonaqueous-stactrolyte secondary battery;

FIG. 2 is a cross sectional view showing enotites conventional nonaqueous-electrolyte secondary battery; FIG. 3 is a cross sectional view showing the structure of a nonsqueous-electrolyte secondary battery according

to the present invention: FIG. 4 is a cross sectional view showing the structure of the nonequeous-electrolyte secondary battery according

FIG. 5 is a perspective view showing a portion including a positive-electrode collector of the nonaqueous-electrolyte secondary battery according to the present invention.

## DESCRIPTION OF THE PREFERRED EMBODIMENTS

[9923] An embodiment of the present invention will now be described with reference to the drawings.

[00247 An embodiment of a nonequeous-electrolyte secondary battery according to the present invention is shown In FIG. 3.

[8925] The nonequeous-electrolyte secondary battery according to this embodiment, so shown in Fig. 3, incorporates a positive electrode 3 having positive-electrode-mix layers 2s and 2b formed on the two sides of a positiveelectrode collector 1. Moreover, a negative electrode 6 incorporates negative-electrode-mb: layers 5a and 5b formed on the two sides of the negative-electrode collector 4. The positive electrode 3 and the negative electrode 6 are wound such that a separator 7 is interposed. The separator 7 is a small-pore film made of polypropylene or polyethylene. Thus, a coil electrode is formed. Thus, insulating members 8 are placed on the two vertical surfaces of the coil electrode, followed by accommodating the coil electrode having the insulating members 8 into a battery can 8.

[0026] A battery cover 10 is joined to the bettery can 9 by crimping the battery cover 10 through a seafing gasket 11. The battery cover 10 and the bettery can 9 are electrically connected to the positive electrods 3 and the negative alectrode 6 respectively through a positive-electrode lead 12 and a negative-electrode lead 13. Thus, the positive electrode and the negative electrode of the battery are formed,

[0027] Note that a current-limiting thin plate 14 serving as a safety unit to provided for the bettery according to this ambodiment. The positive-electrode lead 12 is waitled to the current-limiting thin plate 14 so us to electrically be connected to the battery cover 10 through the current-limiting thin plate 14,

[0028] When the pressure in the battery having the above-mentioned structure has been raised, the current-limiting thin plate 14 is pushed upwards and therefore deformed. Thus, the positive electrode lead 12 is cut such that a portion welded to the current-limiting thin plate 14 is left. As a result, the electric current is limited.

[9929] A cross sectional structure of the nonequeous-electrolyte secondary battery according to the present invention is shown in Fig. 4. In the nonequeous-electrolyte secondary bettery according to this embodiment, the positive-sizetrade-mix layer 2s is, as shown in FIG. 4, formed on only elihor main surface (on the inside) of the positive electrode collector 1 at a position adjacent to an outermost end 3e of the positive electrode 3 antifor an innermost and 3b of the same. No positive-electrods-mix layer is formed on the positive-electrode collector 1 at the outermost end 3s of the positive electrode 3. Moreover, no negative-electrode-mix layer to formed on the negative-electrode collector 4 at an outermost and 6a of the negative electrode 6. That is, only the negative-electrode collector 4 is formed at the outermost end 6a.

[0030] In a direction from the inner portion of the coil electrode 15 to the outer portion of the same, an outermost and of the negative-discinate collector 4, which is the outermost and 6a of the negative electrode 6, is positioned more forwards than the outermost and of the positive-electrods collector 1 which is the outermost end 3a of the positive R sportpele

[9931] As described above, the positive-electrods-mb; layer 2a is formed on only either main surface of the positiveelectrode collector 1 at the position adjacent to the outermost end 3a of the positive electrode 3 and/or the position adjacent to the innermost and 3b of the same. Therefore, the necessity of applying the positive-electrode mix to each

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of the two sides of the positive-electrode collector 1 can be eliminated. Therefore, a portion to which the positiveefectrode-mix layer 2a is applied must be formed on the main surface of the positive-electrode collector 1, as shown In FIG. 5.

[0032] In the nonequeous-electrolyte secondary battery according to the present invention, the negative-electrode laad 13 is formed adjacent to the outermost and of the negative-electrode collector 4 which is the outermost and 6a of the negative electrode 6 of the coal electrode 15. In a direction from the inner portion of the coal electrode 15 to the outer portion, the negative-electrode lead 13 is positioned more forwards than the outermost end of the positive-electrade collector 1 which is the outermost and 3s of the positive electrode 3. Note that the nonaqueous-electrolyte secondary battery according to the present invention has a structure that a positive electrode lead (not shown) of the positive electrode 3 is disposed in the inside portion.

[0033] An assumption will now be described which is made about the following distance of a region in the coll electrode 15 of the nonequeous-electrolyte secundary battery according to the present invention. The distance is a distance in the longitudinal direction from the inner portion of the coll electrode 15 toward the outer portion of the same. The assumption is made that the distance from the outsmost end of the negative-electrode collector 4, which is the outermost and 6a of the negative electrode 6, to the outermost and of the positive-electrode collector 1, which is the outarmost end 3a of the positive electrode 3, to L. Another assumption is made that the diameter of the coll electrode 15 is d. It is preferable that the relationship  $0 \le L \le \pi d$  is satisfied.

[0034] The nonequeous-electrolyte secondary bettery according to the present invention incorporates the coll electrode 15. The coil electrode 15 is formed by laminating the elongated positive electrode 3 and the alongalize negative electrode 6 which are laminated through the separator 7 such that the outermost layer is the positive electrode 3. The positive-electrode-mix layer 2a is formed on only either main surface of the positive-electrode collector 1 at the position adjacent to the autermost and 3s of the positive electrode 3 and/or the position adjacent to the innermost and 3b. In addition, no positive-electrode mix is formed on the positive-electrode collector 1 at the outermost end 3a of the positive electrode 3. Only the positive-electrode collector 1 is formed at the outermost end 3a. Moreover, no negative-electrodemix layer is formed on the negative-electrode collector 4 at the outermost end 6a of the negative electrods 6. Only the negative-electrode collector 4 is formed at the outermost end 6a.

[1035] In the direction from the inner portion of the coll electrode 15 toward the outer portion, the outermost and of the negative electrode collector 4, which is the outermost end 6s of the negative electrods 6. is positioned more forwards than the outermost and of the positive-electrods collector 1 which is the outermost end 3a of the positive electrods 3. Therefore, the quantity of a non-reactive negative-electrode active material in the battery can be reduced. As a result, an effective area can be enlarged correspondingly in the battery. Thus, the inside portion of the battery can effectively be used, causing the energy density to be relised and the lifetime against a cycle operation to be alongated. [0036] The nonequeous-electrolyte secondary battery according to the present invention incorporates the negativeelectrode lead 13 formed adjacent to the outermost end of the negative-electrode collector 4 which is the outermost and 8a of the negative electrode 6. In the direction from the inner portion of the coll electrode 15 lowerd the outer portion of the same, the negative-electrode lead 13 is positioned more forwards than the outermost end of the positivealactrode collector 1 which is the outermost end 3a of the positive electrode 3. Therefore, even if the negative electrode lead 13 plantes the separator 7 disposed between the coll electrode 15 and the battery can 9, the negative-electrode lead 13 is brought into contact with only the battery can 9 which is also the negative electrods. As e result, any internal short circuit occurs, no defect takes place and, therefore, the reliability can be improved.

[0037] In the coll electrode 15 of the nonequeous-electrolyte secondary ballery according to the present invention. the outsimizest end of the negative-electrode collector 4, which is the outsimized and 6e of the negative electrode 6, and the cularmost end of the positive-electrode collector 1, which is the cularmost and 3e of the positive electrode 3, are positioned in the fore-and-off direction from the inner portion of the coll electrode 15 lowerd the outer portion of the same. An assumption is made that the distance from the outermost and of the negative-electrode collector 4 to the outsimost end of the positive-electrode collector 1 is L. Another assumption is made that the dismeter of the coll electrods 15 is d. In this case, it is preferable that the relationship  $0 < L \le \pi d$  is satisfied. If the foregoing structure is employed, no internal short circuit occurs, the energy density is furthermore raised and the illetime against a cycle operation can furthermore be elongated.

[0038] A state will now be considered that the distance L from the outermost end of the negative-electrode collector 4, which is the outermost and 6a of the negative electrode 6, to the outermost and of the positive-electrode collector 1, which is the outermost end 3e of the positive electrode 3, is shorter than the above-mentioned range. In the foregoing case, the outermost end of the positive-electrode collector 1 overlaps the outermost end of the negative-electrode callector 4.

[0039] Therefore, the percent defective is raised though the energy density is not lowered. If the distance L is longer than the above-mentioned range, many portions are produced in which the negotive electrods 6 and the positive electrods 3 are not apposite to each other. Therefore; the energy density is undestrably towered though the persentage

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[8040] The positive electrode 3 and the negative electrode 6 according to the present invention have the abovementioned structures. The mix layers and collectors for constituting the positive electrode 3 and the negative electrode 6 may be known materials.

(0041) The positive-electrode-mix tayers 2a and 2b contain a positive-electrode material, which permits lithium lons to be depended oppositive or an extensive of the conductive material and a binder.

[0042] It is preferable that the positive-electrode material contains LI in a sufficiently large quantity. For example, it is preferable that to employ a composite metal oxide expressed by LIMO<sub>2</sub> (where M is at least one type of a material selected from a group consisting of Co, Ni, Mn, Fe, Ai, V and Ti) and composed of Li and a transition metal; or an interlayer compound containing Li.

[0043] The conductive meterial for imparting conductivity to the positive electrode and the binder for causing the positive-electrode material to be held by the positive-electrode collector may be known materials.

[0044]. The conductive material may be graphite or carbon black, while the binder may be fluorine resin, such as polyvinyildene fluoride.

[0045] The negative-electrode-mix layers 5a and 5b contain the negative-electrode material which permits lithium form to be depet/dedopod and a binder.

[0046] The negative-electrode material may be a carbon material. The carbon material is examplified by pyrocarbon coke (piloh coke, needle coke and petroleum coke), graphite, vitreous carbon, a calcinated organic polymer compound (a material obtained by calcinating phanol resin, furan resin or the filks), carbon filber and active carbon. The negative-electrode material may be cryetalline metal calde or amorphous metal calde which permits lithium lons to be depet/ dedoped, as well as the foregoing carbon material.

[9947] The binder for causing the negative-electrode material to be held by the negative-electrode collector may be a known material. For example, the binder may be fluorine resin, such as polyvinylidens fluoride.

[0048] The battery according to the present invention contains a known horaqueous electrolyte in which an electrolyte is dissolved in nonaqueous solvent, such as organic solvent.

[0049] The organic solvent is not limited particularly. The organic solvent is exemplified by propylene carbonate, ethylene carbonate, 1, 2-dimethoxyethane 1, 2-dimethoxyethane, 1, 2-diathoxyethane, disthyl carbonata, y-butyrolactone, tetrahydrofuran, 1,3-dioxolane, 4-mathyl-1, 3-dioxolane, disthylather, suifotane, methylsuifotane, acetonitrie and propionitrile. Any one of the foregoing material may be employed or a mixture of the came may be employed as mixed solvent.

10050] The electrolyte is not limited particularly. The electrolyte is exemplified by LICIO<sub>4</sub>, LIASF<sub>6</sub>, LIPF<sub>6</sub>, LIBF<sub>4</sub>, LIB (C<sub>6</sub>,H<sub>6</sub>)<sub>4</sub>, LICI, LIBr, LISO<sub>3</sub>CH<sub>3</sub> and LISO<sub>3</sub>CF<sub>3</sub>.

[0061] The material of the separator 7 is not finited perticularly. The material is examplified by woven fabric, unwoven fabric or small-pore film made of synthetic regin, in particular, the small-pore film made of synthetic regin is a preferred material. Moreover, a polyclefine small-pore film is a preferred material in viewpoints of realizing a required thickness, strength of the formed film and resistance of the film. Specifically, the following materials may be employed: a small-pore film made of polycitylene or polypropylene or a small-pore film made of a mature of the foregoing materials.

[0052]. The shape of the collector of the electrode is not limited particularly. A foil structure, a mash structure or a net structure made of expand metal may be employed. It is preferable that the positive-electrode collector 1 is made of, for example, aluminum, stainless steel or nickel, it is preferable that the negative-electrode collector 4 is made of, for example, copper, stainless steel or nickel.

[0050] The battery can 8 may be made of iron, nickel, stainface steel or eluminum. If electrochamical corresion occurs in the nonaqueous electrolyte containing the above-mentioned material during the operation of the battery, platting may be performed.

## 46 Examples

[0054] Examples of the present invention will now be described with results of experiments.

Manufacturing of Samples>

## Sample 1

[0056] mittally, the negative electrode was manufactured as follows.

[0055] Petroleum pitch was employed as a starting material, followed by catcharting the petroleum pitch so that course pitch coke was obtained. The course pitch coke was pulvertized so that powder having an everage particle size of 40 µm was obtained. Then, the obtained powder was calcharted in an inactive gas at 1000°C to remove impurities. Thus, coke powder was obtained.

[9057] Then, 90 parts by weight of thus-obtained coke powder, serving as a carrier for negative-electrode active

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material, and 10 parts by weight of polyvinylidene fluoride (PVDF) serving as a binder were mixed. Thus, a negativeelectrode mix was prepared, followed by disparsing the negative-electrode mix in N-methylpyrolitions which serves as solvent. As a result, negative-electroda mix slurry was obtained. The negative-electrode mix slurry was applied to the two sides of a negative-electrode collector in the form of copper foll having a thickness of 10 µm. Then, the applied solvent was dried. The negative-electrode collector was compression-molded with a roller pressing machine. As a result, an elongated negative electrode was obtained which had a width of 41.5 mm and a length of 250 mm. The thickness of the negative-electrode-mix layer on each side of the negative-electrode collector was 105 µm. The negative-electrode-mix layer was not formed on the negative-electrode collector of the negative electrode at the outermost end. A portion formed by only the negative-electrode collector was created.

[0058] On the other hand, the positive electrode was manufactured as follows.

That is, 0.05 male of lithium carbonate and 1 mole of cabalt carbonate were mixed with each other, and then the relx was calcinated at  $900^{\circ}$ C in air for 5 hours. Thus, LiCoO<sub>2</sub> was obtained.

[0080] Then, obtained LICoO2 was used as a positive-electrode active material such that 91 parts by weight of LICCO<sub>2</sub>, 6 parts by weight of graphile serving as a conductive material and 3 parts by weight of polyvinylidena fluoride (PVDF) serving as a binder were mixed with each other. Thus, a positive-electrode mix was prepared. Then, the obtained positive-electrode mix was dispersed in N-methylpyrolidone so that positive-electrode mix skirry was obtained. The positive-electrode mix sturry was applied to a region of only either aide of a positive-electrode collector made of elongated aluminum foll having a thickness of 20 µm, the region having a length of 247 mm. Then, the positive-electrode mbs sharry was dried. Then, positive-electrode mix sharry was applied to the other main surface of the positive-electrode collector in a region having a length of 167 mm such that a position at which the application was started was made coincide with the above-mentioned positive-electrode mix sturry. Then, the positive-electrode mix sturry was dried, followed by compressing the two sides of the positive-electrode collector with a roll to compression-moid the positiveelectrode collector. Thus, an etongaled positive electrode was obtained which had a width of 39.5 mm. The positive electrode incorporated a portion having two sides on each of which the positive-electrode-mix layor was formed, the portion having a length of 167 mm. A portion of the positive electrode, an either side of which the positive-electrodemix layer was formed, had a length of 80 mm. The thickness of each of the positive-electrode-mix layers was 80 µm. The positive electrode had the outermost end and the innermost end each incorporating a portion in which the positiveelectrode-rnix layer was not formed and in which only the positive-electrode collector was formed.

[0061] The thus-manufactured elongated positive electrode and the negative electrode and two separators, each of which had a thickness of 25 μm and a width of 44 mm and which were in the form of small-pore polypropytene films, ware laminated. The laminate had four layers formed by sequentially laminating the negetive electrode, the separator, the positive electrode and the separator in this sequential order. The laminate was lengthwise wound plural times. Thus, a spiral shape was formed which had a structure that the portion in which only either side of the positive-electrode collector had the positive-electrode-mix layer was first wound and the negative electrode was placed incide. The end of the outermost separator was secured with a tape. Thus, a coll electrode was manufactured. The negative electrode of the coil electrode was longer than the positive electrode of the same. Therefore, in the direction from the inner portion of the coll electrode toward the outer portion, the outermost end of the negative electrode collector which was the outsimps), and of the negative electrode was, as a matter of course, positioned more forwards than the outermost and of the positive electrode collector which was the outermost and of the positive electrode.

[0082] The cuter diameter of the coll collector was 13 mm, while the inner diameter of a hollow portion formed in the central portion of the coll collector was 3.5 mm. The outermost end of the negative-electrode collector which was the outermost end of the negative electrode and the outermost end of the positive-electrode collector which was the outarmost end of the positive electrode were positioned spart from each other in the fore-and-aft direction for distance L which was 35 mm.

[0063] In this embodiment, the negative-electrode lead was positioned at the outermost end of the negative electrode, while the positive-electrode lead was positioned at the innermost and of the positive electrode.

[0064] The thus-manufactured coll electrode was accommodated in an iron battery can applied with nickel plating. An insulating plate was placed on each of the upper and lower sides of the coll electrode. The positive-electrode lead was connected to the battery cover by welding, while the negative-electrode lead was connected to the battery can by

[0065] Than, a nonaqueous electrolyte was prepared by dissolving, at a concentration of 1 molecities, UPFs in a mixed solvent which contained propylene carbonate and diethyl carbonate in the same quantities. Then, 3.0 g of the nonsqueous electrolyte was injected into the bettery can so as to be impregnated into the coil electrode. Than, the battery can was crimped through an insulating sealing gasket applied with asphalt so that the battery cover was secured. Thus, the hermeticity in the battery was maintained.

[0065] Thus, a cylindrical nonequeous-electrolyte secondary battery (having a diameter of 14 mm and a height of 50 mm) was manufactured. The foregoing cylindrical nonequeous-electrolyte secondary battery was called Sample 1 for convenience.

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### Samples 2, 3, 4 to 6, 11 and 12

[0057] Cylindrical nonaqueous-electrolyte secondary batteries were manufectured by a method similar to that for manufacturing Sample 1 except for a structure in which the distance from the outermost end of the negative electrode to the autermost end of the positive electrode which was varied as shown in Table 1. The foregoing secondary betteries were called Samples 4 to 8, 11 and 12. To perform comparisons, cylindrical nonaqueous-electrolyte secondary batteries were manufactured by a method similar to that for manufacturing Sample 1. In this case, as shown in Table 1, the cylindrical nonaqueous-electrolyte secondary batteries were different from Sample 1 as follows: the distance from the cutermost and of the negative electrode and the outermost and of the positive electrode was - 2 mm; the positive electrode collector overlapped the negative-electrode collector in the outermost portion; the distance from the outermost and of the negative electrode to the outsimizationd of the positive electrode was 0 mm; and the end of the negativeelectrode collector and the end of the positive electrode collector coincided with each other in the outermost portion. The foregoing comparative samples were celled Samples 2 and 3 for convenience. Samples 4 to 6, 11, 12, 2 and 3 were structured such that the outer diameter of the coll electrode was made to be 13 mm by adjusting the lengths of the positive-electrode-mix tayers formed on the two sides of the positive electrode and the lengths of the negativeelectrode-mix layers formed on the two sides of the negative electrode.

## Table 1

25	÷	Length of Positive-Electrode- Mix Layer Formed on Either Side of Positive Electrode at Innermost End of Positive Electrode (mm)	Length of Positive-Electrode- Mix Layer Formed on Either Side of Positive Electrode at Outermost End of Positive Electrode (mm)	Length of Positive-Electrode -Mix Layers Formed on Two Sides of Positive Electrode (mm)
_	Sample 1	80	0	167
5 ·	Şample 2	80	0	· 167
	Sample 3	80	0	167
•	Sample 4	60	0	182
	Sample 5	40	0 ·	197
~·· • ·	Sample 6	20	0	201
<b>'</b>	Sample 7	0	5	205
. [	Sample 8	0	15.	202
· l	Sample 9	0	50	176

Sample 10 Sample 11 Sample 12 Sample 13 Sample 14 Sample 15 5 

			<del></del>		
		Length of Negative-Electrode -Mix Layers Formed on Two Sides of Negative Electrode (mm)	Length of Negative-Electrode -Mix Layers Formed on Either Side of Negative Electrode (mm)	Distance from Outermost End of Negative Electrode to Outermost End of Positive Electrode (mm)	
	Sample 1	250	0	35	
	Sample 2	250	0	-2	
	Sample 3	250	0	0	
	Sample 4	245	0	15	
	Sample 5	240	0	10	
	Sample 6	224	0	17	
	Sample 7	213	0	33	
•	Sample 8	220	0	35	
	Sample 9	279	0	50	
	Sample 10	247	0	5	
	Sample 11	243	0	27	
	Sample 12	241	0	38	

Sample 13	239	0 .	43
Sample 14	0	398	35
Sample 15	D	398	0

	Energy Density Ratio (%)	Percent Defective (%)
. Sample 1	100.0	3
5ample 2	100.0	20
Sample 3	100.0	18
Sample 4	. 102.4	. 3
Sample 5	104.8	1 ·
Şample 6	101.9	1 .
Sample 7	100.2	2
Sample 8	101.2	2
Sample 9	97.1	. 1
Sample 10	105.8	5
Sample 11	101.0	1
Sample 12	100.0	2
Sample 13	99.5	3
Sample 14	95.0	2
Sample 15	95.0	20

## 5 <u>Samples 7 to 9</u>

[0068] Processes similar to that for manufacturing Sample 1 were performed except for structures in which the positive-electrode-mix layer was formed on only either side edjecent to the innermost end of the positive electrode. Moreover, the distance from the outermost end of the negative electrode to the outermost end of the positive electrode was varied as shown in Table 1. Thus, cylindrical nonequeous-electrolyte secondary betteries were manufactured. The thus-manufactured cylindrical nonequeous-electrolyte secondary batteries were called Samples 7 to 9 for convenience. Samples 7 to 9 were structured such that the outer diameter of the coll electrode was made to be 13 mm by adjusting the langitive electrode-mix layers formed on the two sides of the positive electrode and the lengths of the negative-electrode-mix layers formed on the two sides of the negative electrode.

### Sample 10

[0069] A cylindrical nonequeous-electrolyte secondary battery was manufactured by a method similar to that for

manufacturing Sample 1 except for structures in which the positive-electrode-mix layer was formed on only either side at positions adjacent to the innermost and outermost ends of the positive electrode. Moreover, the distance from the outermost end of the negative electrode to the outermost end of the positive electrode was varied as shown in Table 1. The thus-manufactured cyfindrical nonaqueous-electrolyte secondary battery was called Sample 10 for convenience. Sample 10 was structured such that the outer diameter of the coll electrode was made to be 13 mm by adjusting the lengths of the positive-electrode-mix layers formed on the two sides of the positive electrode.

## Sample 14 and 15

10 [0070] Cylindrical nonaqueous-electrolyte secondary batteries were manufactured by a method similar to that for manufacturing Sample 1 except for structures in which the positive-electrode-mix layer was formed on the overall length of only either side of the positive electrode and the negative electrode, Moreover, the distance from the outermost end of the negative electrode to the outermost and of the positive electrode was varied as shown in Table 1. The thus-manufactured cylindrical nonaqueous-electrolyte secondary batteries were called Samples 14 and 15 for convenience.
Sample 14 and 15 were structured such that the outer dameter of the coll electrode was made to be 13 mm by adjusting the lengths of the positive-electrode-mix leyers formed on the two sides of the positive-electrode.

#### <Evaluation of Samples>

- [0071] Each of samples 1 to 15 was charged for 8 hours at a charging voltage of 4.20 V with a charging current of 300 mA, and then the foregoing samples were, with a load of 600 mA, discharged to 2.75 V. Thus, the energy densities were measured. Assuming that the result of Sample 3 was 100, the energy densities of the sample betteries with respect to 100 were obtained. Also results were shown in Table 1, Moreover, a percentage defective of each battery was obtained and also results were shown in Table 1.
- 25 (0072) Comparisons were parformed by using results shown in Table 1 among Samples 2, 3 and 15 and samples 1, 4 to 14 according to the present invention. Samples 2, 3 and 15 were structured such that the distance from the outermost end of the negative electrode to the outermost end of the negative electrode collector in the outermost partian. As an afternative to this, the distance from the outermost and of the negative-electrode collector in the outermost end of the positive electrode was 0 mm. Therefore, the end of the negative-electrode collector and the end of the positive-electrode collector and the end of the positive-electrode collector and the end of the positive-electrode collector with each other in the outermost portion. Samples 1, 4 to 14 according to the present invention were manufactured such that the outermost end of the negative-electrode collector which was the outermost end of the negative electrode was, in a direction from the inner portion of the coll electrode lowerd the outer portion of the same; positioned more forwards than the outermost end of the positive-electrode collector which was the outermost end of the positive electrode to the outermost end of the positive electrode collector which was the outermost end of the positive electrode. As a result of the comparisons, the samples according to the present invention enabled the percentage defective to considerably be reduced without reduction in the energy density.
  - [9073] Samples 1 and 4 to 14 according to the present invention were structured such that the outermost end of the negative-electrode collector which was the outermost and of the negative electrode was, in a direction from the inner portion of the coll electrode toward the outer portion of the same, positioned more forwards then the outermost end of the positive-electrode collector which was the outermost end of the positive electrode. Samples 1, 4 to 8 and 10 to 14 among the foregoing samples 1 and 4 to 14 were structured such that the distance L from the outermost end of the negative-electrode collector which was the outermost end of the negative electrode to the outermost end of the positive electrode collector which was the outermost end of the positive electrode collector which was the outermost end of the positive electrode collector which was the outermost end of the positive electrode collector which was the outermost end of the positive electrode collector which was the outermost end of the positive electrode collector which was the outermost end of the positive electrode collector which was the outermost end of the positive electrode collector. When the foregoing electrode of the collector was d. Sample 9 did not satisfy the above mentioned relationship. When the foregoing camples were compared with one another, Sample 9 encountered somewhat reduction in the energy density.
- [0074] Samples 14 and 15, having the electrode mix layer formed on only either side of each of the collectors of both of the positive electrode and the negative electrode, were subjected to a comparison. When the electrode mix layer was formed on only either side of the collector, Sample 14 according to the present invention enabled the parcentage defective to considerably be reduced without reduction in the energy density. On the other hand, Sample 16 to which the present invention is not applied encountered a high percentage defective.
- [0075] As can be understood from the above-manifored results, the structure of the present invention in which the stongated positive and negative electrodes were laminated through separature, followed by winding a laminate such that the positive electrode forms the outermost tayer so that a coll electrode is formed. Moreover, the positive-electrode-mix layer is formed on only either main surface of the collector at a position adjacent to the outermost end of the positive-electrode end/or a position adjacent to the innermost and. At the outermost end of the positive-electrode collector, that is, only the positive-electrode collector is formed. At the outermost and of the negative-electrode collector is not formed on the negative-electrode collector is formed. In the direction from the inner portion of the

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coil electrode toward the outer portion of the same, the outermost end of the negative-electrode collector, which is the outermost end of the negative electrode, is positioned more forwards than the outermost end of the positive-electrode collector which is the outermost and of the positive electrode. Thus, non-reacted ective material for the negative electrode in the battery can be reduced. Thus, an effective area can be enlarged in the battery correspondingly. Therefore, the inside portion of the battery can effectively be used, causing the energy density to be reissed. Thus, elongation of lifetime against cycle operations was confirmed.

[8076] In the present invention, the negative-electrode lead is formed adjacent to the cutermost end of the negative-electrode. In the direction from the inner portion of the coll electrode toward the outer portion of the same, the negative-electrode lead is positioned more forwards than the outermost end of the positive-electrode collector which is the outermost end of the positive electrode. Thus, even the negative-electrode lead be prought that contact with only the battery can which is coll electrode and the battery can, the negative-electrode lead is brought that contact with only the battery can which is the arms negative electrode. Therefore, internal short circuit does not take piece and, therefore, any defect occurs. Thus, the refability can be improved.

[0077] In the present invention, the outermost end of the negative-electrode collector, which is the cutermost end of the negative electrode, and the outermost end of the positive-electrode collector, which is the cutermost end of the positive electrode, are positioned fore-und-aft in the distance from the inner portion of the positive-electrode toward the outer portion of the same. Assuming that the distance from the outermost end of the negative-electrode collector to the outermost end of the positive-electrode collector is L and the distance of the coll electrode is d, the relationship 0 < L < xd is satisfied. Thus, the energy density can furthermore be raised and the lifetime against the cycle operation can furthermore be electrodeed.

[0078] The diameter of the cylindrical nonequeous-electrolyte secondary battery was varied to 18 mm and 20 mm to evaluate each of the manufactured cylindrical nonequeous-electrolyte secondary batteries. Thus, similar tendencies to those shown with the above-mentioned samples were shown.

[0079] An alliptical collectrode was manufactured as the coll electrode so that a nonaqueous-electrolyte secondary battery which was a rectangular battery having a thickness of 9 mm, a width of 34 mm and a height of 48 mm was manufactured so as to be evaluated. The culemost end of the negative-electrode collector which was the outermost end of the negative-electrode was positioned more forwards than the outermost end of the positive-electrode collector which was the outermost and of the positive electrode. The distance from the outermost end of the negative-electrode collector positioned at the outermost end of the positive electrode to the outermost end of the positive-electrode collector positioned at the outermost end of the positive electrode which were positioned fore-end-eft was not longer than the circumference. In this case, reduction of the percentage defective was anabled without reduction in the energy density.

[0080] As described above, the nonequeous-electrolyte secondary battery according to the present Invention Incorporates: a coll electrode formed by laminating an elongated positive electrode and an elongated negative electrode through a separator and by winding a formed taminate such that the positive electrode is positioned at the outermost position, wherein the positive-electrode-mix layer is formed on only either main surface of the collector at the positive-electrode-mix layer is not formed on the positive-electrode-mix layer is not formed on the positive-electrode-mix layer is not formed on the positive-electrode collector at the outermost end of the positive-electrode and only the positive-electrode collector is formed, the negative-electrode-mix layer is not formed on the negative-electrode collector at the outermost end of the negative-electrode collector is formed, and the outermost end of the negative-electrode collector is not end of the negative-electrode collector is not end of the negative-electrode collector in the collector is not end of the negative-electrode collector in the collector is not end of the negative-electrode in the collector is not end of the negative-electrode collector. Therefore, the non-reacted active material for the negative electrode in the battery can be reduced. Thus, the effective bettery area can be enlarged correspondingly. Therefore, the leafle portion of the battery can effectively be used, causing the energy density to be raised and the lifetime against a cycle operation to be elongated.

[0081] The nonequeous-electrolyte secondary battery according to the present invention incorporates the negative-electrode lead formed adjacent to the outermost end of the negative-electrode collector positioned at the outermost end of the negative-electrode collector is, in the direction from the inner portion of the coll electrode toward the outer portion of the same, positioned more forwards than the outermost end of the positive-electrode collector. Thus, even if the negative-electrode lead pierces the separator disposed between the coll electrode the bettery can, the negative-electrode lead is brought into contact with only the battery can which is also the negative electrode. Therefore, internal short circuit can be prevented, any defect can be prevented, and the reliability can be improved.

[0082] The coil electrode of the non-aqueous-electrolyte secondary battery according to the present invention is structured such that the distance it, from the outermost and of the negative-electrode collector positioned at the outermost and of the negative-electrode collector positioned at the unformost and of the positive-electrode collector positioned at the unformost and of the positive electrode which are positioned fore-end-eff in the direction from the inner portion of the coil electrode.

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toward the outer portion of the same satisfies  $0 < L \le \pi d$  on an assumption that the clameter of the coil electrode is d. Thus, internal short circuit can be prevented, the energy density can furthermore be reised and the lifetime against a cycle operation can furthermore be stongated.

[0983] Although the invention has been described in its preferred form and structure with a cartain degree of particularity, it is understood that the present disclusive of the preferred form can be changed in the databs of construction and in the combination and arrangement of parts without departing from the scope of the invention as hereinafter

### Claims

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- A nonaqueous-electrolyte secondary battery comprising:
- e coll electrode formed by laminating an elongated positive electrode which has a positive-electrode-mix layer 15 formed on at least either of main surfaces of a positive-electrode collector and an alongated negative electrode which has a negative-electrode-mix layer formed on at least either main surfaces of a negative-electrode collector and by winding a formed laminate such that seld positive electrode is positioned at the outermost position of said coll electrode, wherein
  - said positive-electrods-mix layer is formed on only either of main surfaces of said collector at the position adjacent to the outermost end of said positive electrode and/or the position adjacent to the innermost end of said positive electrode, said positive electrode-mix layer is not formed on said positive-electrode collector at the outermost end of said positive electrods and only said positive-electrode collector is formed, said negative-electrode-mbi layer is not formed on said negative-electrods collector at the cutermost end of
- said negative electrods and only said negetive-electrode collector is formed, and the outermost end of said negative-electrode collector positioned at the outermost end of said negative electrade is, in the direction from the inner parties of said cell electrode toward the outer parties of said cell electrode, positioned more forwards than the outermost and of said positive-electrode collector.
  - A nonaqueous-electrolyte secondary battery according to claim 1, wherein

said coi) electrode incorporates a negative-electrode lead adjacent to an outermost end of said negativeelectrode collector positioned at the outermost and of eald negative electrode, and said negative-electrode lead is positioned more forwards than the outermost and of said positive-electrode collector positioned at the outermost end of said positive electrode.

A nonequeous-electrolyte secondary battery according to claim 1, wherein

said coll electrode has a structure that distance i. from the outermost and of eald negative-electrode collector positioned at the outermost end of said negative electrode to the outermost end of said positive electrode collector positioned at the outermost end of said positive electrods which are positioned in a fore-end-aft direction from the inner portion of said coil electrode toward the outer portion of said coil electrode sattelles the following relationship on an assumption that the diameter of the coll electrode is ct

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- A nonsqueous-electrolyte secondary battery according to claim 1, wherein sold negative-electrode mix contains a negative-electrode material and a binder.
- 5. A nonaqueous-electrolyte secondary battery according to claim 4, wherein said negative electrods material is at least one type material selected from a group consisting of a crystalline metal colds and an amorphous metal colds which permit doping/dedoping fithlum long.
- 6. A nonequeous-electrolyte secondary battery according to claim 1, wherein said positive-electrode mix contains a positive-electrode material, a conductive material and a binder.
- A nonaqueous-electrolyte secondary textery according to claim B, wherein said positive-electrode material is at least one type material selected from a group consisting of LIMO<sub>2</sub> (where M is at least any one of Co, NJ, Mn, Fe, Al, V and TI) and inturlayer compounds each containing LI.

- A nonaquenus-electrityte secondary battery according to claim 1, wherein
  said separator is made of at least one type material selected from a group consisting of polyethylens and
  polypropylene.
- 9. A nonequeous-electrolyte secondary battery according to claim 1, wherein each positive-electrode collector is made of at least one type material selected from a group consisting of aluminum, stainless steel and nickel.
- 10. A nonaqueous-electrolyte secondary battery according to claim 1, wherein each negative-electrode collector is made of at least one type material selected from a group constalling of copper, stainless steel and nicket.
  - 11. A nonaqueous-electrolyte secondary battery according to claim 1, wherein
- asid nonequeous-electrolyte secondary battery contains a nonequeous electrolyte prepared by dissolving an electrolyte in nonequeous solvent, and seid nonequeous solvent is made of at least one type material selected from a group consisting of propylene carbonate, ethylene carbonate, 1, 2-dimethoxyethane, 1, 2-diethoxyethane, diethylene, diethylene carbonate, 7-butyrolectore, letrahydrofuran, 1,3-dioxolane, 4-methyl-1, 3-dioxolane, diethylether, sulfolane, methylsulfolane, acetonitrile and propionitrile.
  - A nonequeous-electrolyte secondary battery according to claim 11, wherein
     said electrolyte is at least one type material selected from a group consisting of LiClO<sub>4</sub>, LIAsF<sub>6</sub>, LIPF<sub>6</sub>, LIBF<sub>4</sub>.
     LIB(C<sub>6</sub>H<sub>5</sub>)<sub>4</sub>, LICI, LIBI, LISO<sub>3</sub>CH<sub>2</sub> and LISO<sub>3</sub>CF<sub>5</sub>

### Patentensprüche

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1. Sekundärbatlerie mit nicht-wässrigem Elektrolyten, umfassend:

einen Einktrodenwickel erhalten durch Lämfniaren einer längserstreckten positiven Einktrode, bei der auf zumindest einer der Haupfflächen eines positiven Einktrodenkollektors eine Schicht einer positiven Einktrodenmischung aufgeträgen ist, und einer längserstreckten negstiven Einktrode, bei der auf zumindest einer der Haupfflächen eines negstiven Einktrodenkollektors eine negstive Einktrodenmischung eufgebragen ist, wobei des Laminet in der Weise gewickelt ist, dass die positive Einktrode an der äußeren Seite des Einktrodenwickels liegt, wobei

die Schicht der positiven Elektrodenmischung mur an einer Stelle, die en das älutiere Ende der positiven Elektrode angranzt und/oder an einer Stelle, die en das ihnere Ende der positiven Elektrode angranzt, auf eine der Haupflätchen des Kollektors aufgetragen ist, und auf dem äußeren Ende der positiven Elektrode die Schicht der positiven Elektrodenmischung nicht auf den positiven Elektrodenkotiaktor aufgetragen ist, also nur der Kollektor der positiven Elektrode vorliegt,

em äußeren Ende der negetiven Elektrode die Schicht aus der negetiven Elektrodenmischung nicht auf den Kollektor der negetiven Elektrode sufgetragen ist, also nur der Kollektor der negetiven Elektrode verliegt, und wobel in Richtung vom inneren Abschnitt der gewickelten Elektrode zum äußeren Abschnitt der gewickelten Elektrode gesehen das em äußeren Ende der negetiven Elektrode liegende äußere Ende des negetiven Elektrode, trodenkollektors welter voma angecennet ist als das äußere Ende des Kollektors der poeitiven Elektrode.

- 2. Sekundërbattaria mit nicht-wässrigem Elaktrolyten nach Anspruch 1, wobal
- der Elektrodenwickel einen Anschluss für die negetive Elektrode aufweizt, der en des äußere Ende des em äußeren Ende der negativen Elektrode liegenden negativen Elektrodenkollektore enschließt, und der Anschluss der negativen Elektrode weiter vorne liegt als das äußere Ende des em äußeren Ende der positiven Elektrode liegenden positiven Elektrodenkollektore.
- 55 3. Sekund
  ßrbatterte mit nicht-w
  ässrigem Elektrolyten nech Anspruch 1, wobel der Elektrodenwickel an gestaltet ist, dass ein Abstand L zwischen dem 
  äußeren Ende des am 
  äußeren Ende der negativen Elektroden 
  äußeren Ende des am 
  äußeren 
  Ende der negativen Elektrode angeordneten negativen Elektrodenkollektore, wobei diese in L
  ängerichtung

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von dem inneren Abschnitt des Elektrodenwickels zum äußeren Abschnitt des Elektrodenwickels hintereinander angeordnet sind, die folgende Beziehung erfüllt, unter der Annahme, dass der Durchmesser des Elektrodenwickels d anlapdicht

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- Sekundärbatterie mit nicht-w\u00e4serigem Elektrolyten nach Anspruch 1, wobel die negative Elektrodenmischung ein Material f\u00fcr die negative Elektrode und ein Bindemittel umfasst.
- Sekundärbeiterle mit nicht-wässrigem Elektrotyten nach Anspruch 4, wobel
  das Material der negetiven Elektrote zumindest eine Art von Material ist, das ausgewählt ist aus einer Gruppe, welche besteht aus einem kristallinen Metalloxid und einem Amorphen Metalloxid, welche das Dotteren und
  Dedotteren von Lithiumkonen ermöglichen.
- Sekundárbatterie mit nicht-wässingem Eisktrolyten nach Anspruch 1, wobel die positive Elektrodermischung ein Material für die positive Elektrode, ein leitfähiges Material und ein Bindernittel umfasst.
- Sekundärbalteria mit nicht-wässrigem Elektrolyten nech Anspruch 8, wobel
   das positive Elektrodenmaterial zumindest eine Art von Material umfassi, das ausgewählt ist aus einer Gruppe, walche aus LIMO<sub>2</sub> (wobei M zumindest eines ist von Co, NI, Mn, Fe, Al, V und Ti) und Einlagerungsverbindungen besteht, welche jeweils Li enthalten.
- Sekundärbeiterie mit nicht-w\u00e4sstrigern Elektrotyten nach Anspruch 1, wobel
   der Seperatur aus zumindest einem der Materialien herges (eit ist, welche ausgew\u00e4hit sind aus einer Gruppe, bestehend aus Polyethylen und Polypropylen.
  - Sekundärbatterie mit nicht-wässrigem Elektrotyten nach Anspruch 1, wobel
    der positive Elektrodenkollektor aus zumindest einem der Materialien hergestellt ist, die ausgewählt zind aus einer Gruppe, bestehend aus Aluminium, restiretem Stahl und Nickel.
  - Sekundärbatterie mit nicht-wässrigem Elektrolyten nach Anspruch 1, wobei der negative Elektrodenkollekter aus zumindest einem der Materialten hergeeteilt ist, die ausgewählt sind eus einer Gruppe, besiehend aus Kupfer, rostfreiem Stahl und Nitskel.
  - 11. Sekundarbatterle mit nicht-wüssnigern Eiektrolyten nach Anspruch 1, wobei
    - diese einen nicht-wässrigen Elektrotyten enthält, der hergestett ist durch Lösen eines Elektrotyten in einem nicht-wässrigen Lösungsmittel, und das nicht-wässrige Lösungsmittel zumindest aus einem der Materialien hergestellt ist, welche ausgewählt sind aus einer Gruppe, bestehend aus Propytencarbonat, Ethylencarbonat, 1,2-Dimethoxyethen, 1,2-Dimethoxyethen, 1,2-Dimethoxyethen, 1,2-Dimethoxyethen, 1,2-Dimethoxyethen, 1,2-Dimethoxyethen, 1,3-Dioxofen, 4-Methyl-1,3-dioxofen, Digthylether, Suifolan, Methylsuifolan, Acetonitril und Propientitrit.
- 45 12. Sekundárbatterie mit nicht-wässrigem Elaktrotyten nach Anspruch 11, wobel der Elektrotyt zumindest eines der Materialien ist, die ausgewählt sind aus einer Gruppe, bestehend aus LICIO<sub>4</sub>, LIASF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>6</sub>, LIPF<sub>7</sub>, LIPF<sub>8</sub>, LI

## 60 Revendications

- Accumulateur secondaire à électrolyte non equeux comprenant :
- una álactrodo en bobino formás en stratifiant uno étectrodo positivo allongée qui présente una coucha de mélanga d'électrode positive formée sur su moins l'une des gurisces principales d'un collecteur d'électrodo positiva et una électrodo négative allongée qui présente una coucha da mélanga d'électrodo négative formée sur au moins l'une des surfaces principales d'un collecteur d'électrodo négative et en enroutant un stratifié formé de talle sorte que ladite électrode positive est disposée sur la position adérieure de tedite électrode en

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bobine, dans lequel

ladita couche de métange d'électrode postitive est formée sur uniquement l'une des surfaces principales dudit collecteur à la position adjacente de l'extrémité extérieure de ladite électrode positive et/ou à la position adjacente de l'extrémité intérieure de ladité électrode positive, ladile couche de mélange d'électrode positive n'est pas formée sur ledit collocteur d'électrode positive à l'extrémité extérieure de tadite électrode positive et caul ladit collecteur d'électrode positive est formé,

ladite couchs de málange d'électrode négative n'est pas formée sur ladit collecteur d'électrode négative à l'extremité extérieure de ladite électrode négative et soul ledit collecteur d'électrode négative est formé, et l'extrémité extérieure dudit collecteur d'électrode négative disposée à l'extrémité extérieure de ladite électrode négative se trouve, dans la direction allant de la partie interne de ladita électrode en bobine vers la partie externe de lacite électrode en bobine, disposée plus en avant que l'extrémité extérieure dudit collecteur d'électrade positive.

Accumulateur secondaire à électrolyte non equeux selon la revendication 1, dans lequel

ladite électrode en bobino incorpore un conducteur d'électrode négative adjacent à une extrémité extérieure dudit collecteur d'électrode négative disposé à l'extrémité extérieure de latite électrode négative, et ledit conducteur d'électrode négativa est disposé plus en avant de l'extrémité extérieure dudit collecteur d'électrade positive disposé à l'extrémité extérieure de ladite électrade positive.

Accumulateur secondaire à électrolyte non aqueux selon la revendication 1, dans lequel ladite électrode en bobine présente une structure telle que la distance L de l'extrémité extérieure dudit collocteur d'électrode négative disposé à l'extrémité extérieure de ledite électrode négative jusqu'à l'extrémité extérieure dudit collecteur d'électrode positive disposé à l'extrêmité extérieure de ladite électrode positive qui sont disposés dans une direction longitudinale allant de la partie interne de ladita électrode en bobine vers la partie axtame de ladite électrode en bobine satisfait la relation suivante en supposant que le diamètre de l'électrode en bobine est d:

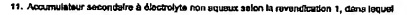
0 < L < xd

- Accumulateur secondaira à électrolyte non aqueux selon la revendication 1, dans lequel ledit mélange d'électroda négative contient un malérieu d'électrode négative et un fiant.
- 5. Accumulatour secondaire à électrolyte non aqueux evion le revendication 4, dans lequel tedit matériau d'électrode négative est un matériau d'eu moins un type choisi parmi un oxyde métallique cristallin et un oxyde matallique amorphe qui permet le dopaga/dédopage d'ions littéurn.
  - Accumulateur secondaire à électrolyte non aqueux selon la revendication 1, dans laquel fedit mélange d'électrode positive confient un matérieu d'électrode positive, un matérieu conducteur et un
  - Accumulateur secondaire à électrolyte non aqueux selon la revendication 6, dans lequel ledit materiau d'électrodo positive est un matériau d'au moine un type cholai parmi LIMO<sub>2</sub> (où M est l'un quelconque parmi Co, Ni, Mn, Fe, Al, V et 11) et des composés de couche intermédiaire contenant chacun Li,
  - Accumulateur secondaire à discirolyte non aqueux seion la revendication 1, dans lequel ledit séparateur est constitué d'un matériau d'au moins un type choisí parmi la polyéthylène et le polypropylène.
  - Accumulateur secondaire à étadrolyte non equeux selon la revendication 1, dans laquet fedit collecteur d'électrode positive est constitué d'un matériau d'au moins un type choisi parmi l'aluminium, l'acier incoydeble et le nickel.
- 10. Accumutateur secondaire à électrolyte non aqueux solon la revendication 1, dans lequel lecti collecteur d'électrode négative est constitué d'un matériau d'au moine un type choisi parmi le cuivre, l'acter inoxydable et la nicical.

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ledit accumulateur secondaire à électrolyte non aqueur contient un électrolyte non aqueux préparé par dissolution d'un électrolyte dans un solvant non equeux, et ledit solvant non aqueux est constitué d'un matériau d'au moins un type choisi parmi le carbonate de propylène, le carbonate d'éthylène, le 1,2-diméthoxyéthane, le 1,2-diéthoxyéthane, le carbonate de diéthyle, la 7-butyrolactone, le tétrahydrofurane, le 1,3-dioxolane, le 4-méthyl-1,3-dioxolane, le diéthyléther, le sulfolane, le méthyleuffolane, l'acétonirile et le propionitrile.

12. Accompliateur secondaire à électrolyte inon equeux selon la revendication 11, dans lequel ledit électrolyte est un matériau d'au moine un type choisi parmi LICIO<sub>4</sub>, LIASF<sub>6</sub>, LIPF<sub>6</sub>, LIBF<sub>4</sub>, LIB(C<sub>5</sub>H<sub>5</sub>)<sub>4</sub>, LICI, LIBC, LISO<sub>3</sub>CH<sub>3</sub> et LISO<sub>3</sub>CF<sub>3</sub>.

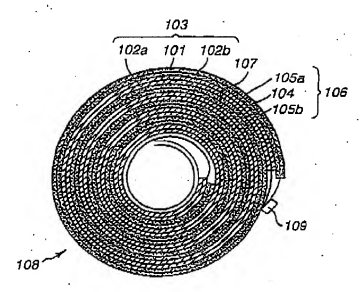


FIG.1

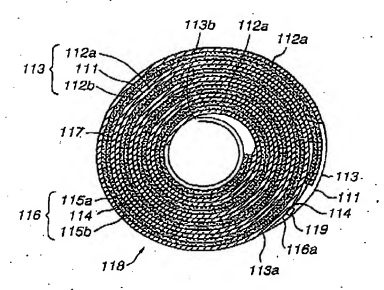


FIG.2

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QUALLION LEGAL

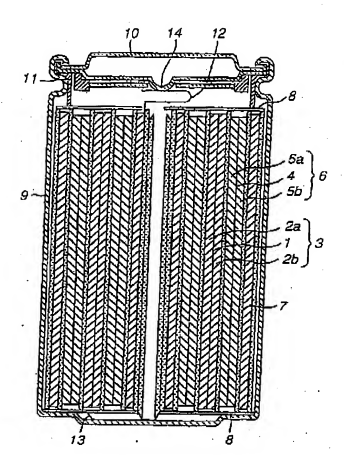


FIG.3

QUALLION LEGAL

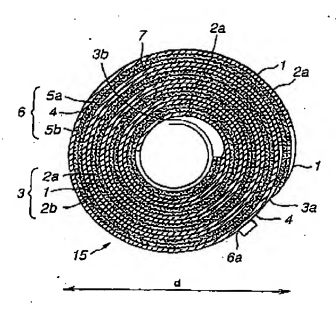


FIG.4

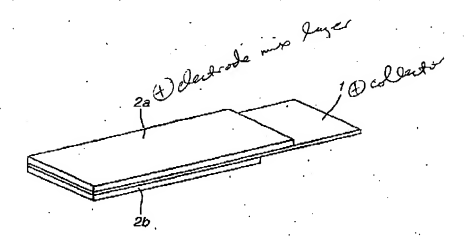


FIG.5